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10	Annex F	
11	ICCVAM/NICEATM BG1Luc4E2 ER TA – Antagonist Protocol	

13

November 8, 2010

1601 E. Geer St., Suite S

Durham, NC 2770413

08 November 2010

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165	LIST OF	ACRONYMS AND ABBREVIATIONS
166	13 mm test tube	13 x 100 mm glass test tubes
167	DMEM	Dulbecco's Modification of Eagle's Medium
168	DMSO	Dimethyl Sulfoxide
169 170	DMSO Control	1% v/v dilution of DMSO in tissue culture media used as a vehicle control
171	E2	17β-estradiol
172	E2 Control	$2.5 \times 10^{-5} \mu g/mL$ E2 used as a control.
173 174	IC ₅₀ Value	Concentration that produces a half-maximal response as calculated using the four parameter Hill function.
175	ER	Estrogen Receptor
176 177 178	Estrogen-free DMEM	DMEM (phenol red free), supplemented with 1 % Penicillin/Streptomycin, 2 % L-Glutamine, and 5% Charcoaldextran treated FBS
179	FBS	Fetal Bovine Serum
180 181	Flavone/E2 Control	25 μg/mL flavone + 2.5 x 10 ⁻⁵ μg/mL E2, used as a weak positive control.
182	G418	Gentamycin
183 184	Ral/E2 Reference Standard	Nine point dilution of raloxifene HCl + $2.5 \times 10^{-5} 17\beta$ -estradiol reference standard for the LUMI-CELL® ER antagonist assay
185	RPMI	RPMI 1640 growth medium
186	TA	Transcriptional Activation
187	T25	25 cm ² tissue culture flask
188	T75	75 cm² tissue culture flask
189	T150	150 cm ² tissue culture flask
190		

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226	1.0	PURPOSE
440	1.0	LUNIUSI

- This protocol is designed to evaluate coded test substances for potential estrogen receptor (ER) antagonist
- 228 activity using the LUMI-CELL® ER assay.
- 229 **2.0 SPONSOR**
- 230 The National Toxicology Program (NTP) Interagency Center for the Evaluation of Alternative
- Toxicological Methods (NICEATM), P.O. Box 12233 Research Triangle Park, NC 27709
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280	Chemistr	y Resources Group Leader		
281	National	Institute of Environmental Health Sciences		
282		06, P.O. Box 12233		
283		Triangle Park, NC 27709		
284	Phone: 9	19-541-3473		
285	3.0	DEFINITIONS		
286		• Dosing Solution: The test substance, control substance, or reference standard solution		
287		which is to be placed into the tissue culture wells for experimentation.		
288		• Raw Data: Raw data includes information that has been collected but not formatted or		
289		analyzed, and consists of the following:		
290		 Data recorded in the Study Notebook 		
291		 Computer printout of initial luminometer data 		
292		 Other data collected as part of GLP compliance, e.g.: 		
293		 Equipment logs and calibration records 		
294		 Test substance and tissue culture media preparation logs 		
295		 Cryogenic freezer inventory logs 		
296		• Soluble: Test substance exists in a clear solution without visible cloudiness or		
297		precipitate.		
298		• Study Notebook: The study notebook contains recordings of all activities related to the		
299		conduct of the BG1LUC4E2 ER TA TA antagonist assay.		
300		• Test Substances: Substances supplied to the testing laboratories that are coded and		
301		distributed such that only the Project Officer, Study Management Team (SMT), and the		

302 303 304		Substance Inventory and Distribution Management have knowledge of their true identity. The test substances will be purchased, aliquoted, coded, and distributed by the Supplier under the guidance of the NIEHS/NTP Project Officer and the SMT.
305	4.0	TESTING FACILITY AND KEY PERSONNEL ¹
306	4.1	Testing Facility
307	Xenobio	otic Detection Systems, Inc. (XDS), 1601 E. Geer St., Durham, NC 27704
308	4.2	Key Personnel
309		• Study Director: John Gordon, Ph.D.
310		• Quality Assurance Director: Mr. Carlos Daniel
311	5.0	IDENTIFICATION OF TEST AND CONTROL SUBSTANCES
312	5.1	Test Substances
313 314		stances are coded and will be provided to participating laboratories by the Substance Inventory ribution Management team.
315	5.2	Controls
316	Controls	s for the ER antagonist protocol are as follows:
317 318		control (dimethyl sulfoxide [DMSO]): 1% v/v dilution of DMSO (CASRN 67-68-5) diluted in ulture media.
319	Ral/E2 r	reference standard for range finder testing: Three concentrations (1.56 x 10 ⁻³ ,
320 321		0^{-4} , and 9.77 x 10^{-5} μg/mL) of raloxifene HCl (Ral), CASRN 84449-90-1, plus a fixed ration (2.5 x 10^{-5} μg/mL) of 17β-estradiol (E2), CASRN: 50-28-2, in duplicate wells.
322 323		reference standard for comprehensive testing: A serial dilution of Ral plus a fixed concentration $0^{-5} \mu g/mL$) of E2 consisting of nine concentrations of Ral/E2 in duplicate wells.
324 325	E2 control.	<i>rol</i> : 17β-estradiol, 2.5 x 10^{-5} µg/mL E2 in tissue culture media used as a base line negative
326 327		/E2 Control: Flavone, CASRN 525-82-6, 25 μg/mL, with 2.5 x 10 ⁻⁵ μg/mL E2 in tissue culture sed as a weak positive control.
328	6.0	OVERVIEW OF GENERAL PROCEDURES FOR ANTAGONIST TESTING
329 330 331	•	erimental procedures are to be carried out under aseptic conditions and all solutions, glassware, ware, pipettes, etc., shall be sterile. All methods and procedures shall be documented in the study k.

¹ Testing facility and personnel information are provided as an example.

Antagonist range finder testing is conducted on 96-well plates using three concentrations of Ral/E2 (1.56

 $x = 10^{-3}$, 3.91 x 10^{-4} , and 9.77 x 10^{-5} µg/mL Ral) with 2.50 x 10^{-5} µg/mL E2) in duplicate as the reference

standard, with three replicate wells for the E2 and DMSO controls.

Comprehensive testing is conducted on 96-well plates using nine concentrations of Ral/E2 in duplicate as

the reference standard (**Table 6-1**). Four replicate wells for the DMSO control, Flavone/E2 and E2

controls are included on each plate.

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Table 6-1 Concentrations of Ral/E2 Reference Standard Used for Comprehensive Testing

Raloxifene Concentrations ¹	E2 Concentrations
1.25 x 10 ⁻²	2.5 x 10 ⁻⁵
6.25 x 10 ⁻³	2.5 x 10 ⁻⁵
3.13×10^{-3}	2.5 x 10 ⁻⁵
1.56 x 10 ⁻³	2.5 x 10 ⁻⁵
7.81 x 10 ⁻⁴	2.5 x 10 ⁻⁵
3.91 x 10 ⁻⁴	2.5 x 10 ⁻⁵
1.95 x 10 ⁻⁴	2.5 x 10 ⁻⁵
9.77 x 10 ⁻⁵	2.5 x 10 ⁻⁵
4.88 x 10 ⁻⁵	2.5 x 10 ⁻⁵

¹Concentrations are presented in μg/mL.

Visual observations for cell viability are conducted for all experimental plates just prior to BG1LUC4E2

ER TA evaluation, as outlined in **Section 11.4**.

Luminescence data, measured in relative light units (RLUs), is corrected for background luminescence by subtracting the mean RLU value of the vehicle control (DMSO) wells from the RLU measurements for each of the other wells of the 96-well plate. Data is then transferred into Excel® data management spreadsheets and GraphPad PRISM® 4.0 statistical software, graphed, and evaluated for a positive or negative response as follows:

- A response is considered positive for antagonist activity when the average adjusted RLU for a given concentration is less than the mean RLU value minus three times the standard deviation for the E2 control.
- Any luminescence at or above this threshold is considered a negative response.

For substances that are positive at one or more concentrations, the concentration of test substance that causes a half-maximal response (the relative IC_{50}) is calculated using a Hill function analysis. The Hill function is a four-parameter logistic mathematical model relating the substance concentration to the response (typically following a sigmoidal curve) using the equation below

$$Y = Bottom + \frac{Top - Bottom}{1 + 10^{(logIC50-X)HillSlope}}$$

where Y = response (i.e., relative light units); X = the logarithm of concentration; Bottom = the minimum response; Top = the maximum response; log IC_{50} = the logarithm of X as the response midway between

- Top and Bottom; and HillSlope describes the steepness of the curve. The model calculates the best fit for
- 359 the Top, Bottom, HillSlope, and IC₅₀ parameters. See **Section 13.6.5** for more details.
- 360 Acceptance or rejection of a test is based on evaluation of reference standard and control results from
- ach experiment conducted on a 96-well plate. Results for these controls are compared to historical results
- compiled in the historical database, as seen in **Section 16.0**.

6.1 Range Finder Testing

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- Antagonist range finding for coded substances consists of a seven-point 1:10 serial dilution using
- duplicate wells per concentration. Concentrations for comprehensive testing are selected based on the
- response observed in range finder testing. If necessary, a second range finder test can be conducted to
- clarify the optimal concentration range to test (see Section 14.0).

6.2 Comprehensive Testing

- 369 Comprehensive antagonist testing for coded substances consists of 11-point serial dilutions, with each
- 370 concentration tested in triplicate wells of the 96-well plate. Three separate experiments are conducted for
- 371 comprehensive testing on three separate days, except during Phases III and IV of the validation effort, in
- which comprehensive testing experiments are conducted once (see Section 15.0).

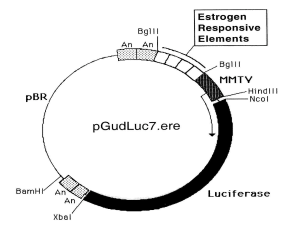
373 7.0 MATERIALS FOR BG1LUC4E2 ER TA ANTAGONIST TESTING

- This section provides the materials needed to conduct BG1LUC4E2 ER TA testing, with associated brand
- names/vendors² in brackets.

376 7.1 **BG1Luc4E2** Cells:

- Human ovarian cancer cell line stably transfected with a plasmid containing an estrogen response element
- 378 (Figure 7-1) [XDS].

379 Figure 7-1 pGudLuc7.ERE Plasmid.



²Brand names and vendors should not be considered an endorsement by the U.S. Government or any member of the U.S. Government; such information is provided as examples.

381	7.2	Technical Equipment:
382 383 384		cal equipment may be obtained from Fisher Scientific International, Inc. (Liberty Lane NH, USA 03842). Equivalent technical equipment from another commercial source can be
385		• Analytical balance (Cat. No. 01-910-320)
386 387		 Berthold Orion 1 Microplate Luminometer [Berthold CatNo.: Orion 1 MPL3] or equivalent and dedicated computer
388		• Biological safety hood, class II, and stand (Cat. No. 16-108-99)
389 390		• Centrifuge (low speed, tabletop with swinging bucket rotor) (Cat. No. 04-978-50 centrifuge, and 05-103B rotor)
391		• Combustion test kit (CO ₂ monitoring) (Cat. No. 10-884-1)
392		• Drummond diaphragm pipetter (Cat. No. 13-681-15)
393		• Freezers, -20°C (Cat. No. 13-986-150), and -70°C (Cat. No. 13-990-86)
394		• Hand tally counter (Cat. No. 07905-6)
395		• Hemocytometer, cell counter (Cat. No. 02-671-5)
396		• Light microscope, inverted (Cat. No. 12-561-INV)
397		• Light microscope, upright (Cat. No. 12-561-3M)
398		• Liquid nitrogen flask (Cat. No. 11-675-92)
399		• Micropipetter, repeating (Cat. No. 21-380-9)
400		• Pipetters, air displacement, single channel (0.5 –10ml (Cat. No. 21-377-191), 2 –20 ml
401 402		(Cat. No. 21-377-287), 20 – 200 ml (Cat. No. 21-377-298), 200 - 1000 ml (Cat. No. 21-377-195))
403		• Refrigerator/freezer (Cat. No. 13-986-106A)
404		• Shaker for 96-well plates (Cat. No. 14-271-9)
405		• Sodium hydroxide (Cat. No. 5318-500)
406		• Sonicating water bath (Cat. No. 15-335-30)
407		• Tissue culture incubator with CO ₂ and temperature control (Cat. No. 11-689-4)
408		• Vacuum pump with liquid trap (side arm Erlenmeyer) (Cat. No. 01-092-29)
409		• Vortex mixer (Cat. No. 12-814)
410	Equipmen	t should be maintained and calibrated as per GLP guidelines and individual laboratory SOPs.
411	7.3	Reference Standard, Controls, and Tissue Culture Supplies
412 413 414	dates. Tiss	culture reagents must be labeled to indicate source, identity, storage conditions and expiration sue culture solutions must be labeled to indicate concentration, stability (where known), and in and expiration dates.

Equivalent tissue culture media and sera from another commercial source can be used, but must first be 415 416 tested as described in **Section 17.0** to determine suitability for use in this test method. 417 The following are the necessary tissue culture reagents and possible sources based on their use in the pre-418 validation studies: 419 BackSeal-96/384, white adhesive bottom seal for 96-well and 384-well microplate 420 [Perkin-Elmer, Cat. No. 6005199] 421 17 β-estradiol (CAS RN: 50-28-2) [Sigma-Aldrich, Cat. No. E8875] 422 CellTiter-Glo[®] Luminescent Cell Viability Assay [Promega Cat. No. G7572] 423 Cryovial, 2 mL (Corning Costar) [Fisher Scientific Cat. No. 03-374-21] 424 Culture tube 13 x 100mm (case) [Thomas Scientific Cat. No.: 10009186R38]³ 425 Culture tube, 50 mL conical (Corning Costar) [Fisher Scientific Cat. No. 05-526C] 426 DMSO, U.S.P. analytical grade. [Sigma-Aldrich, Cat. No. 34869-100ML] 427 Dulbecco's Modification of Eagle's Medium (DMEM), containing 4.5 g/L glucose, with 428 sodium pyruvate, without phenol red or L-glutamine [Mediatech/Cellgro, Cat. No. 17-429 205-CV] 430 Fetal Bovine Serum [Mediatech/Cellgro Cat. No. MT 35-010-CV] 431 Fetal Bovine Serum, charcoal/dextran treated, triple 0.1 µm sterile filtered [Hyclone, Cat. 432 No. SH30068.03] 433 Flavone (CASRN: 525-82-6) [Sigma-Aldrich, Cat. No. F2003] 434 Gentamycin Sulfate (G418), 50 mg/mL [Mediatech/Cellgro Cat. No. 30-234-CR] 435 L-glutamine, 29.2 mg/mL [Cellgro, Cat. No. 25005-CI] 436 Luciferase Assay System (10-Pack) [Promega Cat. No. E1501] 437 Lysis Solution 5X [Promega, Cat. No. E1531] 438 Penicillin/streptomycin solution, 5000 I.U. penicillin, 5000 μg/mL streptomycin [Cellgro, 439 Cat. No. 30-001-CI]. 440 Phosphate buffered saline (PBS, 1X) without calcium and magnesium [Cellgro, Cat. No. 441 21-040-CV] 442 Pipettes, serological: 2.0 mL [Sigma-Aldrich, Cat. No. P1736], 5.0 mL [Sigma-Aldrich, 443 Cat. No. P1986], 25 mL [Sigma-Aldrich, Cat. No. P2486] 444 Raloxifene (CASRN 84449-90-1) [Sigma-Aldrich Cat. No. R1402] 445 RPMI 1640 medium, containing L-glutamine [Mediatech, Cat. No. 10-040-CV] Tissue culture flasks (Corning-Costar): 25 cm² (T25) [Fisher Cat. No. 10-126-28]; 75 cm² 446 447 (T75) [Fisher Cat. No. 10-126-37]; and 150 cm² (T150) [Fisher Cat. No. 10-126-34]

³If glass tubes can not be obtained from Thomas Scientific, the preference is for flint glass, then lime glass, then borosilicate glass.

448 Tissue culture plates (Corning-Costar): 96-well [Thomas Scientific Cat. No. 6916A05] 449 Trypsin (10X), 2.5% in Hank's balanced salt solution (HBSS), without calcium and 450 magnesium, without phenol red [Cellgro, Cat. No. 25-054-CI]. 451 All reagent lot numbers and expiration dates must be recorded in the study notebook. 452 8.0 PREPARATION OF TISSUE CULTURE MEDIA AND SOLUTIONS 453 All tissue culture media and media supplements must be quality tested before use in experiments (see 454 **Section 15.0**). 455 **RPMI 1640 Growth Medium (RPMI)** 8.1 456 RPMI 1640 is supplemented with 0.9% Pen-Strep and 8.0% FBS to make RPMI growth medium (RPMI). 457 Procedure for one 549 mL bottle: 458 Remove FBS from -70°C freezer, and Pen-Strep from -20°C freezer and allow to 459 equilibrate to room temperature. 460 2. Add 44 mL of FBS and 5 mL Pen-Strep to the bottle of RPMI 1640. 461 3. Label RPMI bottle as indicated in **Section 7.3** 462 Store at 2-8°C for no longer than six months or until the shortest expiration date of any media 463 component. 464 8.2 **Estrogen-Free DMEM Medium** 465 DMEM is supplemented to contain 4.5% charcoal/dextran treated FBS, 1.9% L-glutamine, 0.9% Pen-466 Strep. 467 Procedure for one 539 mL bottle: 468 Remove charcoal/dextran treated FBS from -70°C freezer, and L-glutamine and Pen-469 Strep from -20°C freezer and allow to equilibrate to room temperature. 470 Add 24 mL of charcoal/dextran treated FBS, 10 mL L-glutamine, and 5 mL Pen-Strep to 471 one 500 mL bottle of DMEM. 472 Label estrogen-free DMEM bottle as indicated in Section 7.3 473 Store at 2-8 °C for no longer than six months or until the shortest expiration date of any media 474 component. 475 8.3 **1X Trypsin Solution** 476 1X Trypsin solution is prepared by dilution from a 10X premixed stock solution. The 10X stock solution 477 should be stored in 10 mL aliquots in a -20°C freezer. 478 Procedure for making 100 mL of 1X trypsin: 479 Remove a 10mL aliquot of 10X trypsin from -20°C freezer and allow to equilibrate to 480 room temperature.

Aliquot 1 mL Trypsin (10X) along with 9 mL of 1X PBS into ten 15 mL centrifuge tubes.

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482		3. Label 1X trypsin aliquots as indicated in Section 7.3
483	1X Trypsin	should be stored at -20°C.
484	8.4	1X Lysis Solution
485 486		tion is prepared by dilution from a 5X premixed stock solution. Both the 5X and 1X solutions eatedly freeze-thawed.
487	The proceed	dure for making 10 mL of 1X lysis solution:
488		1. Thaw the 5X Promega Lysis solution and allow it to reach room temperature.
489		2. Remove 2 mL of 5X solution and place it in a 15 mL conical centrifuge tube.
490		3. Add 8 mL of distilled, de-ionized water to the conical tube.
491		4. Cap and shake gently until solutions are mixed.
492	Store at -2	$0^{\circ}C$ for no longer than 1 year from receipt.
493	8.5	Reconstituted Luciferase Reagent
494	Luciferase	reagent consists of two components, luciferase buffer and lyophilized luciferase substrate.
495	For long-te	erm storage, unopened containers of the luciferase buffer and lyophilized luciferase substrate
496	can be stor	red at -70°C for up to six months.
497	To reconst	itute luciferase reagent:
498 499		1. Remove luciferase buffer and luciferase substrate from -70°C freezer and allow them to equilibrate to room temperature.
500		2. Add 10 mL of luciferase buffer solution to luciferase substrate container and swirl or
501		vortex to mix, the Luciferase substrate should readily go into solution.
502		3. Luciferase substrate should readily go into solution.
503		4. After solutions are mixed aliquot to a 15mL centrifuge tube.
504		5. Store complete solution at –20°C.
505	Reconstitu	ted luciferase reagent is stable for 1 month at -20°C.
506 507	9.0	OVERVIEW OF PROPAGATION AND EXPERIMENTAL PLATING OF BG1Luc4E2 CELLS
508 509 510 511 512 513	monolayer humidity, a under an ir must be no	uc4E2 (BG-1) cells are stored in liquid nitrogen in 2 mL cryovials. BG-1 cells are grown as a in tissue culture flasks in a dedicated tissue culture incubator at $37^{\circ}\text{C} \pm 1^{\circ}\text{C}$, $90\% \pm 5\%$ and $5.0\% \pm 1\%$ CO ₂ /air. The cells should be examined on a daily basis during working days everted phase contrast microscope, and any changes in morphology and adhesive properties sted in the study notebook. flasks containing cells at 80% to 90% confluence will usually yield a sufficient number of cells
514		e 96-well plates for use in experiments.

Procedures for Thawing Cells and Establishing Tissue Cultures

- Warm all tissue culture media and solutions to room temperature by placing them under the tissue culture
- 517 hood several hours before use.
- All tissue culture media, media supplements, and tissue culture plasticware must be quality tested before
- use in experiments (Section 17.0).

520 9.1.1 Thawing Cells

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- 1. Remove a cryovial of frozen BG-1 cells from the liquid nitrogen flask.
- 522 2. Facilitate rapid thawing by loosening the top slightly (do not remove top) to release trapped gasses and retightening it. Roll vial between palms.
- 3. Use a micropipette to transfer cells to a 50 mL conical centrifuge tube.
 - 4. Rinse cryovial twice with 1X PBS and add PBS rinse material to the conical tube.
- 5. Add 20 mL of RPMI to the conical tube.
- 527 6. Centrifuge at 1000 x g for eight min. If a pellet of cells has not formed, centrifuge for an additional 5 minutes.
 - 7. Aspirate media from pellet and re-suspend it in 5 mL RPMI, drawing the pellet repeatedly through a 1.0 mL serological pipette to break up any clumps of cells.
 - 8. Transfer cells to a T25 flask, place them in an incubator (see conditions in **Section 9.0**) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).

533 9.1.2 <u>Establishing Tissue Cultures</u>

- Once cells have reached 80% to 90% confluence, transfer the cells to a T75 flask by performing, for example, the following steps:
 - 1. Remove the T25 flask from the incubator.
 - 2. Aspirate the RPMI, then add 5 mL 1X PBS, making sure that the cells are coated with PBS.
 - 3. Aspirate 1X PBS, then add 1 to 2 mL 1X trypsin to the T25 flask, gently swirling the flask to coat all cells with the trypsin.
 - 4. Place the flask in an incubator (see conditions in **Section 9.0**) for 5 to 10 min.
 - 5. Detach cells by hitting the side of the flask sharply against the palm or heel of the hand.
 - 6. Confirm cell detachment by examination under an inverted microscope. If cells have not detached, return the flask to the incubator for an additional 2 minutes, then hit the flask again.
 - 7. After cells have detached, add 5 mL PBS, and transfer the suspended cells to a 50 mL centrifuge tube. Wash the flask one additional time with 5 mL PBS.
 - 8. Immediately add 20 mL RPMI to the conical tube to inhibit further cellular digestion by residual trypsin.
 - 9. Pellet the cells by centrifugation, as described in **Section 9.1.1**, and re-suspend the cells in 10 mL RPMI medium.

552 553	10.	Draw the pellet repeatedly through a 25 mL serological pipette to break up clumps of cells
554 555	11.	Transfer cells to a T75 flask, then place the flask in an incubator (see conditions in Section 9.0) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).
556 557	When cells have example, the fo	re reached 80% to 90% confluency, transfer them into a T150 flask by performing, for
558	•	Remove the T75 flask from the incubator, aspirate the old media and add 5 mL 1X PBS.
559 560		Aspirate 1X PBS, add 2 mL of 1X trypsin to the flask, and place it in an incubator (see conditions in Section 9.0) for 5 to 10 min.
561	14.	Repeat steps 5 through 11 in Section 9.1.2 , re-suspending the pellet in 20 mL of RPMI.
562 563	15.	Transfer cells to a T150 flask and place it in the incubator (see conditions in Section 9.0) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).
564	16.	Remove the T150 flask from the incubator.
565	17.	Aspirate the RPMI and add 5 mL 1X PBS.
566 567	18.	Aspirate 1X PBS and add 3 mL 1X trypsin to the T150 flask, making sure that the cells are coated with the trypsin.
568	19.	Incubate cells in an incubator (see conditions in Section 9.0) for 5 to 10 min.
569	20.	Detach cells by hitting the side of the flask sharply against the palm or heel of the hand.
570 571 572	21.	Confirm cell detachment by examination under an inverted microscope. If cells have not detached, return the flask to the incubator for an additional 2 minutes, then hit the flask again.
573 574 575	22.	
576 577	23.	Immediately add 20 mL RPMI to the conical tube to inhibit further cellular digestion by residual trypsin.
578 579	24.	Centrifuge at 1000 x g for eight minutes. If a pellet of cells has not formed, centrifuge for an additional 5 minutes.
580 581	25.	Aspirate the media from the pellet and re-suspend it in 40 mL RPMI, drawing the pellet repeatedly through a 25 mL serological pipette to break up any clumps of cells.
582 583 584	26.	Transfer 20 mL of cell suspension to each of two T150 flasks, place them in an incubator (see conditions in Section 9.0) and grow to 80% to 90% confluence (approximately 48 to 72 hrs).
585 586		going Tissue Culture Maintenance, Conditioning in Estrogen-free Medium, and ting Cells for Experimentation
587 588	•	procedure is used to condition the BG1Luc4E2 cells to an estrogen-free environment prior ells in 96-well plates for analysis of estrogen dependent induction of luciferase activity.

589 To start the tissue culture maintenance and estrogen-free conditioning, split the two T150 culture flasks 590 into four T150 flasks. Two of these flasks will be used for continuing tissue culture and will use the 591 RPMI media mentioned above. The other two flasks will be cultured in estrogen-free DMEM for 592 experimental use. Extra care must be taken to avoid contaminating the estrogen-free cells with RPMI. 593 Remove both T150 flasks from the incubator. 594 2. Aspirate the medium and rinse the cells with 5 mL 1X PBS. 595 Aspirate 1X PBS, then add 3 mL 1X trypsin to the flasks, gently swirling the flask to coat 596 all cells with the trypsin. 597 Incubate cells in the incubator (see conditions in **Section 9.0**) for 5 to 10 min. 4. 598 5. Detach cells by hitting the side of the flask sharply against the palm or heel of the hand. 599 Confirm cell detachment by examination under an inverted microscope. If cells have not 600 detached, return the flask to the incubator for an additional 2 minutes, then hit the flask 601 again. 602 7. After cells have detached, add 5 mL 1X PBS to the first T150 flask and transfer the 603 suspended cells to the second T150 flask. 604 Transfer the contents of both flasks to a 50 mL conical tube. Repeat step 7 with an 605 additional 5 mL 1X PBS and transfer to the 50 mL conical tube. 606 Immediately add 20 mL estrogen-free DMEM to the 50 mL conical tube to inhibit further 607 cellular digestion by residual trypsin. 608 10. Centrifuge at 1000 x g for eight minutes. If a pellet of cells has not formed, centrifuge for 609 an additional 5 minutes. 610 11. Aspirate media from pellet and re-suspend it in 4 mL estrogen-free DMEM, drawing the 611 pellet repeatedly through a 1 mL serological pipette to break up clumps of cells. 612 At this point, cells are ready to be divided into the ongoing tissue culture and estrogen-free conditioning 613 groups. 614 9.2.1 Ongoing Tissue Culture Maintenance 615 Add 20 mL RPMI to two T150 flasks. 616 Add 220 µL G418 to the RPMI in the T150 flasks 2. 617 3. Add 1 mL of cell suspension from **Section 9.2 step 11** to each flask. 618 Place T150 flasks in tissue culture incubator (see conditions in **Section 9.0**) and grow to 619 80% to 90% confluence (approximately 48 to 72 hrs). 620 Tissue culture medium may need to be changed 24 hours after addition of G418 to 621 remove cells that have died because they do not express reporter plasmid. 622 G418 does not need to be added to the flasks a second time. 6.

624 9.2.2 Conditioning in Estrogen-free Medium

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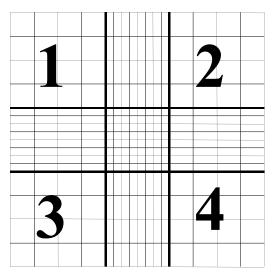
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Repeat Section 9.2 steps 1-11 for ongoing tissue culture maintenance.

625 Add 20 mL estrogen-free DMEM to two T150 flasks. 626 2. Add 150 µL G418 to the estrogen-free DMEM in the T150 flasks. 627 3. Add 1 mL of cell suspension from Section 9.2 step 11 to each flask. 628 Tissue culture medium may need to be changed 24 hours after addition of G418 to 4. 629 remove cells that have died because they do not express reporter plasmid. 630 5. G418 does not need to be added to the flasks a second time. 631 6. Place the T150 flasks in the incubator (see conditions in **Section 9.0**) and grow to 80% to 632 90% confluence (approximately 48 to 72 hrs). 633 9.2.3 Plating Cells Grown in Estrogen-free DMEM for Experimentation 634 Remove the T150 flasks that have been conditioned in estrogen-free DMEM for 48 to 72 635 hours from the incubator. 636 Aspirate the medium, then rinse the cells with 5 mL 1X PBS. 2. 637 Aspirate 1X PBS, then add 3 mL 1X trypsin to the flasks, gently swirling the flask to coat 3 638 all cells with the trypsin. 639 Place the flasks in an incubator (see conditions in **Section 9.0**) for 5 to 10 min. 4. 640 5. Detach cells by hitting the side of the flask sharply against the palm or the heel of the 641 hand. 642 Confirm cell detachment by examination under an inverted microscope. If cells have not 643 detached, return the flask to the incubator for 2 additional minutes, then hit the flask 644 again. 645 7. After cells have detached, add 5 mL 1X PBS and transfer the suspended cells from the 646 T150 flask to a 50 mL conical tube. Add an additional 5 mL PBS to the flask, then 647 transfer to the 50 mL conical tube. 648 Immediately add 20 mL estrogen-free DMEM to each conical tube to inhibit further 649 cellular digestion by residual trypsin. 650 9. Centrifuge at 1000 x g for eight minutes. If a pellet of cells has not formed, centrifuge for 651 an additional 5 minutes. 652 10. Aspirate off the media from the pellet and re-suspend it in 20 mL DMEM, drawing the 653 pellet repeatedly through a 25 mL serological pipette to break up any clumps of cells. 654 11. Pipette 15 µL of the cell suspension into the "v" shaped slot on the hemocytometer. 655 Ensure that the solution covers the entire surface area of the hemocytometer grid, and 656 allow cells to settle before counting. 657 12. Using 100x magnification, view the counting grid. 658 13. The counting grid on the hemocytometer consists of nine sections, four of which are 659 counted (upper left, upper right, lower left, and lower right, see Figure 9-1). Each section 660 counted consists of four by four grids. Starting at the top left and moving clockwise, 661 count all cells in each of the four by four grids. Some cells will be touching the outside

borders of the square, but only count those that touch the top and right borders of the square. This value is then used in the calculation below to get to the desired concentration of 200,000 cells/mL.

Figure 9-1 Hemocytometer Counting Grid.



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The volume of each square is 10⁻⁴ mL, therefore:

Cells/mL = (average number per grid) x 10^{-4} mL. x 1/(starting dilution).

Starting dilution: 20mL (for T150 flasks)

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Harvested cells for a T150 flask are suspended in 20 mL of estrogen-free DMEM and sampled for determination of concentration of cells/mL.

- Example Calculation:
- Grids 1, 2, 3, and 4 are counted and provide the following data:
- 674 0 50, 51, 49, and 50: average number of cells per grid is equal to 50.
- Cells/mL = 50 cells per grid \div 10⁻⁴ mL volume of grid = 50 X 10⁻⁴ cells/mL (or 500,000 cells/mL)
- Total # of Cells Harvested = 500,000 cells/mL x 20 mL
- Desired Concentration (or Concentration Final) = 200,000 cells/mL
- Formula: (Concentration $_{Final}$ x Volume $_{Final}$ = Concentration $_{Initial}$ x Volume $_{Initial}$)
- 679 Concentration Final = 200,000 cells/mL
- 680 Concentration Initial = 500,000 cells/mL
- Volume $_{Initial} = 20 \text{ mL}$
- Volume $_{Final}$ to be solved for.
- Therefore: 200,000 cells/mL x Volume $_{Final}$ = 500,000 cells/mL x 20 mL
- Solving for Volume $_{Final}$ we find = 50 mL

- Therefore, add 30 mL of estrogen-free DMEM to the cell suspension for a total volume of 50 mL, which will yield the desired concentration of 200,000 cells/mL for plating.
 - 14. This dilution scheme will give a concentration of 200,000 cells/mL. 200 mL of this cell suspension is used for each well of a 96-well plate (i.e., 40,000 cells per well).
 - 15. Remove a 96-well plate from its sterile packaging. Use a repeater pipetter to pipette 200 μ L of cell suspension into each well to be used for the testing of coded substances, reference standard and controls (**note**: add 200 μ L of estrogen-free DMEM only to any wells not being used for testing).
 - 16. Incubate plate(s) in an incubator (see conditions in **Section 9.0**) for a minimum of 24 hours, but no longer than 48 hours before dosing.
- Two T150 flasks containing cells at 80% to 90% confluence will typically yield sufficient cells to fill four 96-well plates (not including the perimeter wells).

697 10.0 PREPARATION OF TEST SUBSTANCES

- The solvent used for dissolution of test substances is 100% DMSO. All test substances should be allowed
- 699 to equilibrate to room temperature before being dissolved and diluted. Test substance solutions (except
- for reference standards and controls) should not be prepared in bulk for use in subsequent tests. Test
- substances are to be used within 24 hours of preparation. Solutions should not have noticeable precipitate
- or cloudiness.

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- All information on weighing, solubility testing, and calculation of final concentrations for test substances,
- reference standards and controls is to be recorded in the study notebook.

705 10.1 Determination of Test Substance Solubility

- 1. Prepare a 200 mg/mL solution of the test substance in 100% DMSO in a 4 mL conical tube.
- 2. Vortex to mix.
 - 3. If the test substance does not dissolve at 200 mg/mL, prepare a 20 mg/mL solution and vortex as above.
 - 4. If the test substance does not dissolve at 20 mg/mL solution, prepare a 2 mg/mL solution in a 4 mL conical tube and vortex as above.
 - 5. If the test substance does not dissolve at 2 mg/mL, prepare a 0.2 mg/mL solution in a 4 mL conical tube and vortex as above.
 - 6. Continue testing, using 1/10 less substance in each subsequent attempt until test substance is solubilized in DMSO.
- Once the test substance has fully dissolved in 100% DMSO, the test substance is ready to be used for
- 718 BG1LUC4E2 ER TA testing.
- The Testing Facility shall forward the results from the solubility tests assay to the SMT through the
- designated contacts in electronic format and hard copy upon completion of testing.

- 721 11.0 PREPARATION OF REFERENCE STANDARD, CONTROL AND TEST
- 722 SUBSTANCE STOCK SOLUTIONS FOR RANGE FINDER AND COMPREHENSIVE
- 723 TESTING
- All information on preparation of test substances, reference standards and controls is to be recorded in the
- 325 study notebook.
- 726 11.1 Preparation of Ral/E2 Stock Solutions
- E2 and raloxifene stocks are prepared separately and then combined into Ral/E2 stocks, which are then
- used to prepare dosing solutions in **Section 12**.
- 729 11.1.1 E2 Stock Solution
- The final concentration of the E2 stock solution is 5.0 x 10⁻³ µg/mL. Prepare the E2 stock as shown in
- 731 **Table 11-1**.

732 Table 11-1 Preparation of E2 Stock Solution

Step #	Action	DMSO	E2 Concentration
1	Make a 10 mg/mL stock solution in 100% DMSO in a 4mL vial.	-	10 mg/mL
2	Transfer 10 μL E2 solution from Step #1 to a	Add 990 μL of 100%	100 μg/mL
	new 4 mL vial.	DMSO. Vortex to mix.	100 μg/IIIL
2	Transfer 10 μL E2 solution from Step #2 to a	Add 990 μL of 100%	1 μg/mL
3	new 4mL vial.	DMSO. Vortex to mix.	1 μg/IIIL
4	Transfer 100 μL E2 solution from Step #3 to a	Add 9.90 mL of 100%	1.0 x 10 ⁻² μg/mL
4	new glass container large enough to hold 15 mL.	DMSO. Vortex to mix.	1.0 x 10 μg/IIIL
5	Transfer 5 mL E2 solution from Step #4 to a	Add 5 mL of 100%	5.0 x 10 ⁻³ μg/mL
3	new glass container large enough to hold 15 mL	DMSO. Vortex to mix.	J.O X 10 µg/IIIL

- 733 11.1.2 Raloxifene Stock Solution
- 734 Prepare a 2.5 μg/mL raloxifene working stock solution as shown in **Table 11-2**.

735 Table 11-2 Preparation of Raloxifene Stock Solution

Step #	Action	DMSO	Raloxifene Concentration
1	Make a 10 mg/mL solution of raloxifene in a 4 mL glass vial.	-	$1.0 \times 10^4 \mu g/mL$
2	Transfer 10 μL raloxifene solution from Step #1 to a new 4 mL vial.	Add 990 μL of 100% DMSO. Vortex to mix.	100 μg/mL
3	Transfer 150 μL raloxifene solution from Step #2 to a new 4 mL vial.	Add 2.850 mL of 100% DMSO. Vortex to mix.	5 μg/mL
4	Transfer 1.5 mL raloxifene solution from Step #3 to a new 13 mm test tube.	Add 1.5 mL of 100% DMSO. Vortex to mix.	2.5 μg/mL

736 11.2 Ral/E2 Range Finder Testing Stock

- 737 11.2.1 Raloxifene Dilutions
- Number three 4 mL vials with the numbers 1 to 3 and use the raloxifene solution prepared in **Section**
- 739 **11.1.2** to make raloxifene dilutions as shown **Table 11-3**.

740 Table 11-3 Preparation of Raloxifene Dilutions for Range Finder Testing

Step #	Action	DMSO	Raloxifene Concentration
1	Transfer 250 μL of the 2.5 μg/mL raloxifene working	Add 750 μL of 100% DMSO	6.25 x 10 ⁻¹ μg/mL
1	stock solution to a 4 mL tube	and vortex	0.23 x 10 μg/IIIL
2	Transfer 500 μL of the 6.25 x 10 ⁻¹ μg/mL raloxifene	Add 500 μL of 100% DMSO	3.13 x 10 ⁻¹ μg/mL
2	solution to a 4 mL tube	and vortex	3.13 x 10 μg/IIIL
2	Transfer 250 μL of the 3.13 x 10 ⁻¹ μg/mL raloxifene	Add 750 μL of 100% DMSO	7.81 x 10 ⁻² μg/mL
3	solution to a 4 mL tube	and vortex	7.81 x 10 µg/IIIL
4	Transfer 125 μL of the 7.81 x 10 ⁻² μg/mL raloxifene	Add 375 μL of 100% DMSO	1.95 x 10 ⁻² μg/mL
4	solution to a 4 mL tube	and vortex	1.93 x 10 µg/IIIL

- 741 11.2.2 <u>Preparation of Ral/E2 Range Finder Working Stocks:</u>
- Through 3 and add 500 μL of the 5 x 10⁻³ mg/mL E2
- solution prepared in **Section 11.1.1** to each tube. Add 500 μ L of the 3.13 x 10⁻¹, 7.81 x 10⁻², and 1.95 x
- 744 10⁻² μg/mL raloxifene solutions prepared in **Section 11.2.1** to tubes 1, 2, and 3 respectively. Vortex each
- tube to mix. The final concentrations for raloxifene and E2 are listed in **Table 11-4**.

746 Table 11-4 Concentrations of Raloxifene and E2 in the Ral/E2 Range Finder Working Stocks

Tube #	Raloxifene (mg/ml)	E2 (mg/ml)
1	1.56 x 10 ⁻¹	2.5 x 10 ⁻³
2	3.91 x 10 ⁻²	2.5 x 10 ⁻³
3	9.77 x 10 ⁻³	2.5 x 10 ⁻³

747 11.3 Ral/E2 Comprehensive Testing Stock

748 11.3.1 Raloxifene Dilutions

Use the raloxifene solution prepared in **Section 11.1.2** to make a nine-point serial dilution of raloxifene as

750 shown **Table 11-5**.

751 Table 11-5 Preparation of Raloxifene Dilutions for Comprehensive Testing

Step #	Action	DMSO	Discard	Raloxifene Concentration
1	Transfer 500 µL of the raloxifene working stock solution to a new 4 mL vial.	-	-	2.5 μg/mL
2	Transfer 500 µL of the raloxifene working stock solution to a new 4 mL vial.	Add 500 μL of 100% DMSO. Vortex to mix.	-	1.25 μg/mL
3	Transfer 500 µL raloxifene solution from Step #2 to a new 4 mL vial.	Add 500 μL of 100% DMSO. Vortex to mix.	-	6.25 x 10 ⁻¹ μg/mL
4	Transfer 500 µL raloxifene solution from Step #3 to a new 4 mL vial.	Add 500 μL of 100% DMSO. Vortex to mix.	-	3.13 x 10 ⁻¹ μg/mL
5	Transfer 500 µL raloxifene solution from Step #4 to a new 4 mL vial.	Add 500 µL of 100% DMSO. Vortex to mix.	-	1.56 x 10 ⁻¹ μg/mL
6	Transfer 500 µL raloxifene solution from Step #5 to a new 4 mL vial.	Add 500 µL of 100% DMSO. Vortex to mix.	-	7.81 x 10 ⁻² μg/mL
7	Transfer 500 µL raloxifene solution from Step #6 to a new 4 mL vial.	Add 500 µL of 100% DMSO. Vortex to mix.	-	3.91 x 10 ⁻² μg/mL
8	Transfer 500 µL raloxifene solution from Step #7 to a new 4 mL vial.	Add 500 µL of 100% DMSO. Vortex to mix.		1.95 x 10 ⁻² μg/mL
9	Transfer 500 μL raloxifene solution from Step #8 to a new 4 mL vial.	Add 500 μL of 100% DMSO. Vortex to mix.	Discard 500 μL from Tube #9	9.77 x 10 ⁻³ μg/mL

752 11.3.2Preparation of Ral/E2 Comprehensive Testing Working Stocks:

- 753 Add 500 μ L of the 5 x 10⁻³ mg/mL E2 solution prepared in **Section 11.1.1** to each of the 9 raloxifene
- dilution vials (including the working stock solution in Tube #1). Vortex each tube to mix. The final
- concentrations for raloxifene and E2 are listed in **Table 11-6**.

Table 11-6 Concentrations of Raloxifene and E2 in the Ral/E2 Working Stocks

Tube #	Raloxifene (mg/mL)	E2 (mg/mL)
1	1.25	2.5 x 10 ⁻³
2	6.25 x 10 ⁻¹	2.5 x 10 ⁻³
3	3.13×10^{1}	2.5 x 10 ⁻³
4	1.56 x 10 ⁻¹	2.5 x 10 ⁻³
5	7.81×10^2	2.5 x 10 ⁻³
6	3.91 x 10 ⁻²	2.5 x 10 ⁻³
7	1.95 x 10 ⁻²	2.5 x 10 ⁻³
8	9.77 x 10 ⁻³	2.5 x 10 ⁻³
9	4.88 x 10 ⁻³	2.5 x 10 ⁻³

757 11.4 Flavone/E2 Stock Solution

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- 758 To prepare the flavone/E2 stock solution, proceed as follows:
- 759 1. Prepare 1 mL of 5 mg/mL flavone
- 760 2. Add 1 mL of 5x10⁻³ mg/mL E2 (prepared as in **Section 11.1.1**) to the 10 mg/mL flavone. 761 This will make a working solution of 2.5 mg/mL flavone with 2.5x10⁻³ mg/mL E2.

762 12.0 PREPARATION OF REFERENCE STANDARD, CONTROL AND TEST 763 SUBSTANCE DOSING SOLUTIONS FOR RANGE FINDER AND 764 COMPREHENSIVE TESTING

765 12.1 Preparation of Reference Standard and Control Dosing Solutions - Range Finder Testing

- Range finder testing is conducted on 96-well plates using three concentrations of Ral/E2 in duplicate as
- the reference standard. Three replicate wells for the DMSO, and E2 controls are included on each plate.
- All "dosing solutions" of test substance concentrations are to be expressed as µg/mL in the study
- notebook and in all laboratory reports. Dosing solutions are to be used within 24 hours of preparation.

770 12.1.1 <u>Preparation of Ral/E2 Reference Standard Range Finder Dosing Solutions</u>

- 1. Label three 13 mm glass tubes with the numbers 1 to 3.
- 2. Add 6 µL Ral/E2 stock from tube #1 (Section 11.2.2) to 13 mm glass test tube #1.
- 3. Add 6 μ L of Ral/E2 stock from tube #2 from **Section 11.2.2** to the 13 mm glass test tube labeled #2. Repeat for tube #3.
- 4. Add 600 μL of estrogen-free DMEM to each tube and vortex.

776 12.1.2 Preparation of DMSO Control Range Finder Dosing Solution

- 1. Add 8 μL of 100% DMSO to a 13 mm glass test tube.
- 778 2. Add 800 μL of estrogen-free DMEM to each tube and vortex.

779 12.1.3 Preparation of E2 Control Range Finder Dosing Solution

- 780 1. Add 4 μL of the E2 stock from **Section 11.1.1** to a 13 mm glass test tube.
- 781 2. Add 4 μ L of 100% DMSO to the tube.
- 782 3. Add $800 \mu L$ of estrogen-free DMEM to the tube and vortex to mix.

783 12.2 Preparation of Test Substance Dosing Solutions for Range Finder Testing

Range finder experiments are used to determine the concentrations of test substance to be used during comprehensive testing. Antagonist range finding for coded substances consists of seven-point 1:10 serial dilutions in duplicate.

To prepare test substance dosing solutions:

 Label two sets of seven glass 13 mm test tubes with the numbers 1 through 7 and place them in a test tube rack. Perform a serial dilution of test substance as shown in **Table 12-**1 using one set of tubes.

Table 12-1 Preparation of Test Substance Serial Dilution for Range Finder Testing

Tube #	100% DMSO	Test Substance ¹	Final Volume
1	-	100 μL of test substance solution from Section 10.1	100 μL
2	90 μL	10 μL of test substance solution from Section 10.1	100 μL
3	90 μL	10 μL from Tube #2	100 μL
4	90 μL	10 μL from Tube #3	100 μL
5	90 μL	10 μL from Tube #4	100 μL
6	90 μL	10 μL from Tube #5	100 μL
7	90 μL	10 μL from Tube #6	100 μL

Vortex tubes #2 through 6 before removing test substance/DMSO solution to place in the next tube in the series.

2. Transfer test substance/DMSO solutions to the second set of labeled tubes and add E2 as shown in **Table 12-2**.

Table 12-2 Addition of E2 to Test Substance Serial Dilution for Range Finder Testing

Tube Number	Test Substance	E2	Estrogen-free DMEM ³	Final Volume
1	Transfer 4 μL of test substance from Tube #1 in Section 12.2 step 1 to a new tube	Add 4 μL of the 5 x 10 ⁻³ μg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL
2	Transfer 4 µL of test substance from Tube #2 to a new tube	Add 4 μL of the 5 x 10 ⁻³ μg/mL E2 solution prepared in Section 11.1.1 Vortex to mix.	800 μL	808 μL
3	Transfer 4 µL of test substance from Tube #3 to a new tube	Add 4 μL of the 5 x 10 ⁻³ μg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL
4	Transfer 4 µL of test substance from Tube #4 to a new tube	Add 4 µL of the 5 x 10 ⁻³ µg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL
5	Transfer 4 µL of test substance from Tube #5 to a new tube	Add 4 µL of the 5 x 10 ⁻³ µg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL
6	Transfer 4 µL of test substance from Tube #6 to a new tube	Add 4 µL of the 5 x 10 ⁻³ µg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL
7	Transfer 4 µL of test substance from Tube #7 to a new tube	Add 4 µL of the 5 x 10 ⁻³ µg/mL E2 solution prepared in Section 11.1.1 . Vortex to mix.	800 μL	808 μL

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Determination of whether a substance is positive in range finder testing and selection of starting concentrations for comprehensive testing will be discussed in **Section 14.0**.

12.3 Preparation of Reference Standard and Control Dosing Solutions for Comprehensive Testing

Comprehensive testing is conducted on 96-well plates using nine concentrations of Ral/E2 in duplicate as the reference standard. Four replicate wells for the DMSO, E2 and flavone/E2 controls are included on each plate.

All "dosing solutions" of test substance concentrations are to be expressed as $\mu g/mL$ in the study notebook and in all laboratory reports.

Store dosing solutions at room temperature. Use within 24 hours of preparation.

12.3.1 <u>Preparation of Ral/E2 Reference Standard Dosing Solutions for Comprehensive Testing</u> In preparation for making Ral/E2 1:2 serial dilutions, label two sets of nine glass 13 mm test tubes with the numbers 1 through 9 and place them in a test tube rack. Tube number 1 will contain the highest concentration of raloxifene (**Table 12-3**).

Table 12-3 Preparation of Ral/E2 Reference Standard Dosing Solution for Comprehensive Testing

Tube Number	Ral/E2 Stock	Estrogen- free DMEM	Final Volume
1	6 μL of Tube #1 from Section 11.3.2	600 μL	606 μL
2	6 μL of Tube #2 from Section 11.3.2	600 μL	606 μL
3	6 μL of Tube #3 from Section 11.3.2	600 μL	606 μL
4	6 μL of Tube #4 from Section 11.3.2	600 μL	606 μL
5	6 μL of Tube #5 from Section 11.3.2	600 μL	606 μL
6	6 μL of Tube #6 from Section 11.3.2	600 μL	606 μL
7	6 μL of Tube #7 from Section 11.3.2	600 μL	606 μL
8	6 μL of Tube #8 from Section 11.3.2	600 μL	606 μL
9	6 μL of Tube #9 from Section 11.3.2	600 μL	606 μL

Preparation of DMSO Control Comprehensive Testing Dosing Solution

- 1. Add 10 µL of 100% DMSO to a 13 mm glass test tube.
- 2. Add 1000 μL of estrogen-free DMEM to the tube and vortex to mix.

816 12.3.3 Preparation of E2 Control Comprehensive Testing Dosing Solution

- 1. Add 5 μL of the E2 stock from **Section 11.1.1** to a 13 mm glass test tube.
- 818 2. Add 5 μ L of 100% DMSO to the tube.
- 3. Add 1000 µL of estrogen-free DMEM to the tube and vortex to mix.

820 12.3.4 Preparation of Flavone/E2 Control Comprehensive Dosing Solution

- 1. Add 10 μL of flavone/E2 from **Section 11.4** to a 13 mm glass test tube.
- 822 2. Add 1000 μL of estrogen-free DMEM to the tube and vortex to mix.

823 12.4 Preparation of Test Substance Dosing Solutions for Comprehensive Testing

- 824 Comprehensive testing experiments are used to determine whether a substance possesses ER antagonist
- activity in the BG1LUC4E2 ER TA test method. Antagonist comprehensive testing for coded substances
- consists of either an 11-point 1:2 serial dilution, or an 11-point 1:5 serial dilution with each concentration
- tested in triplicate wells of the 96-well plate.
- 828 12.4.1 Preparation of Test Substance 1:2 Serial Dilutions for
- 829 *Comprehensive Testing*

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- Start the 11-point serial dilution according to criteria in **Section 14.0**.
- To make test substance 1:2 serial dilutions for comprehensive testing:
- 1. label eleven 4 mL conical tubes with numbers 1 through 11 and place them in a tube rack
 - 2. label eleven 13 mm glass test tubes with numbers 1 through 11, place them in a tube rack and add 800 μL of estrogen-free DMEM to each tube
- Prepare dilution of test substance as shown in **Table 12-4**.

Table 12-4 Preparation of Test Substance 1:2 Serial Dilutions for Comprehensive Testing

Tube Number	100% DMSO	Test Substance ¹	Discard	E2 Testing Stock	Estrogen- free DMEM ²	Final Volume
1	1	4 μL of test substance solution from Section 10.2.4 step 1	-	4 μL	800 μL	808 μL
2	4 μL	4 μL of test substance solution from Section 10.2.4 step 1	-	4 μL	800 μL	808 μL
3	4 μL	4 μL from Tube #2	-	4 μL	800 μL	808 μL
4	4 μL	4 μL from Tube #3	-	4 μL	800 μL	808 μL
5	4 μL	4 μL from Tube #4	-	4 μL	800 μL	808 μL
6	4 μL	4 μL from Tube #5	-	4 μL	800 μL	808 μL
7	4 μL	4 μL from Tube #6	-	4 μL	800 μL	808 μL
8	4 μL	4 μL from Tube #7	-	4 μL	800 μL	808 μL
9	4 μL	4 μL from Tube #8	-	4 μL	800 μL	808 μL
10	4 μL	4 μL from Tube #9	-	4 μL	800 μL	808 μL
11	4 μL	4 μL from Tube #10	4 μL	4 μL	800 μL	808 μL

Nortex tubes #2 through 10 before removing test substance/DMSO solution to place in the next tube in the series.

12.4.2 Preparation of Test Substance 1:5 Serial Dilutions for

Comprehensive Testing

- Start the 11-point serial dilution according to criteria in **Section 14.0**.
- To make test substance 1:5 serial dilutions for comprehensive testing:
 - 1. label eleven 4 mL conical tubes with numbers 1 through 11 and place them in a tube rack
 - 2. label eleven 13 mm glass test tubes with numbers 1 through 11, place them in a tube rack and add 800 μ L of estrogen-free DMEM to each tube
- Prepare dilution of test substance as shown in **Table 12-5**.

^{838 &}lt;sup>2</sup>Vortex all tubes to mix media, test substance, and E2.

847 Table 12-5 Preparation of Test Substance 1:5 Dilutions for Comprehensive Testing

Tube Number	100% DMSO	Test Substance ¹	Discard	E2 Testing Stock	Estrogen-free DMEM ²	Final Volume
1	-	4 μL of test substance solution from Section 10.2.4 step 1	-	4 μL	800 μL	808 μL
2	16 μL	4 μL of test substance solution from Section 10.2.4 step 1	-	4 μL	800 μL	808 μL
3	16 μL	4 μL from Tube #2	-	4 μL	800 μL	808 μL
4	16 μL	4 μL from Tube #3	-	4 μL	800 μL	808 μL
5	16 μL	4 μL from Tube #4	-	4 μL	800 μL	808 μL
6	16 μL	4 μL from Tube #5	-	4 μL	800 μL	808 μL
7	16 μL	4 μL from Tube #6	-	4 μL	800 μL	808 μL
8	16 μL	4 μL from Tube #7	-	4 μL	800 μL	808 μL
9	16 μL	4 μL from Tube #8	-	4 μL	800 μL	808 μL
10	16 μL	4 μL from Tube #9	-	4 μL	800 μL	808 μL
11	16 μL	4 μL from Tube #10	20 μL	4 μL	800 μL	808 μL

Vortex tubes #2 through 10 before removing test substance/DMSO solution to place in the next tube in the series.

13.0 GENERAL PROCEDURES FOR THE TESTING OF CODED SUBSTANCES

Range finder experiments are used to determine the concentrations of test substance to be used during comprehensive testing. Comprehensive testing experiments are used to determine whether a substance

possesses ER antagonist activity in the BG1LUC4E2 ER TA test method.

854 General procedures for range finder and comprehensive testing are nearly identical. For specific details

(such as plate layout) of range finder testing see Section 14.0. For specific details of comprehensive

856 testing, see Section 15.0.

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13.1 Application of Reference Standard, Control and Test Substances

- 1. Remove the 96-well plates (from **Section 9.2.3 step 18)** from the incubator; inspect them using an inverted microscope. Only use plates in which the cells in all wells receive a score of 1 according to **Table 11-1**.
- 2. Remove medium by inverting the plate onto blotter paper. Gently tap plate against the bench surface to remove residual liquid trapped in the wells.
- 3. Add 200 µL of medium, reference standard, control or test substance to each well (see **Sections 14.0** and **15.0** for specific plate layouts).
- 4. Return plates to incubator (see **Section 9.0** for details) for 19 to 24 hours to allow maximal induction of luciferase activity in the cells.

13.1.1 Preparation of Excel® Data Analysis Template For Range Finder Testing

- 1. In Excel®, open a new "AntRFTemplate" and save it with the appropriate project name as indicated in the NICEATM Style Guide.
- 2. Fill out the table at the top of the "Raw Data" worksheet with information regarding the Microplate reader used, Reading Direction, No. of Intervals, Tot. Meas. Time/Well (s),

²Vortex all tubes to mix media, test substance, and E2.

872			etc. (note: this information can be permanently added to the default template
873			"AntRFTemplate" on a laboratory specific basis).
874		3.	Add the following information regarding the assay to the "Compound Tracking"
875			worksheet.
876			 Plate # - Enter the experiment ID or plate number into cell E1
877			 Cell Lot # - Enter the passage or lot number of the cells used for this
878			experiment into cell B5
879 880			■ DMSO and Media Lot #'s – Enter the lot numbers for the DMSO and Media in cells B6 and B7
881			 Test Substance Code – Enter the test substance codes into cells C14 to C19
882			Name: Enter the experimenter name into cell G6
883			■ Date: Enter the experiment date in the format day\month\year into cell G10
884 885			Comments: - Enter any comments about the experiment in this box (e.g., plate
		4	contaminated)
886		4.	Enter the following substance testing information to the "List" worksheet:
887 888			 Concentration – Type in the test substance concentration in μg/ml in descending order.
889 890			Any specific comments about the test substance or condition of the wells should be entered into this sheet, in the comments section
891 892			 All of the remaining cells on the "List" worksheet should populate automatically.
893 894 895			The "Template", "Compound Mixing" and "Visual Inspection" worksheet should automatically populate with the information entered into the "Compound Tracking" and "List" worksheet.
896		5.	Save the newly named project file.
897		6.	Print out either the "List" or "Template" worksheet for help with dosing the 96-well
898			plate. Sign and date the print out and store in study notebook.
899	13.1.2	Pre	eparation of Excel® Data Analysis Template for Comprehensive Testing
900		1.	In Excel®, open a new "AntCTTemplate" and save it with the appropriate project name as
901			indicated in the NICEATM Style Guide.
902		2.	Fill out the table at the top of the "Raw Data" worksheet with information regarding the
903			Microplate reader used, Reading Direction, No. of Intervals, Tot. Meas. Time/Well (s),
904			etc. (note: this information can be permanently added to the default template
905			"AntCTTemplate" on a laboratory specific basis).
906		3.	On the "Compound Tracking" worksheet, enter the following information:
907			 Plate # - Enter the experiment ID or plate number into cell E1

908 909			 Cell Lot # - Enter the passage or lot number of the cells used for this experiment into cell C5
910 911			 DMSO and Media Lot #'s – Enter the lot numbers for the DMSO and Media in cells C6 and C7
912 913			 Test Substance Code – Enter the test substance codes into cells C15 and C16. Enter the test substance dilution into cells D15 and D16.
914			 Name: Enter the experimenter name into cell F6
915			 Date: Enter the experiment date in the format day\month\year into cell G10
916 917			 Comments: - Enter any comments about the experiment in this box (e.g., plate contaminated)
918		4.	Enter the following substance testing information to the "List" worksheet:
919 920			 Concentration – Type in the test substance concentration in μg/ml in descending order.
921 922			 Any specific comments about the test substance or condition of the wells should be entered into this sheet, in the comments section
923 924			 All of the remaining cells on the "List" worksheet should populate automatically.
925 926 927			• The "Template", "Compound Mixing" and "Visual Inspection" worksheet should automatically populate with the information entered into the "Compound Tracking" and "List" worksheet.
928		5.	Save the newly named project file.
929 930		6.	Print out either the "List" or "Template" worksheet for help with dosing the 96-well plate. Sign and date the print out and store in study notebook.
931	13.2	Vis	ual Evaluation of Cell Viability
932 933 934		1.	19 to 24 hours after dosing the plate, remove the plate from the incubator and remove the media from the wells by inverting the plate onto blotter paper. Gently tap plate against the bench surface to remove residual liquid trapped in the wells.
935 936		2.	Use a repeat pipetter to add 50 μL 1X PBS to all wells. Immediately remove PBS by inversion.
937 938		3.	Using an inverted microscope, inspect all of the wells used in the 96-well plate and record the visual observations using the scores in Table 13-1 .
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939 Table 13-1 Visual Observation Scoring

Viability Score	Brief Description ¹
1	Normal Cell Morphology and Cell Density
2	Altered Cell Morphology and/or Small Gaps between Cells
3	Altered Cell Morphology and/or Large Gaps between Cells
4	Few (or no) Visible Cells
P	Wells containing precipitation are to be noted with "P"

P40 Reference photomicrographs are provided in the BG1LUC4E2 ER TA Validation Study "Visual Observation Cell Viability Manual."

942 13.3 Lysis of Cells for BG1LUC4E2 ER TA

- 1. Apply the reflective white backing tape to the bottom of the 96-well plate (this will increase the effectiveness of the luminometer).
- 2. Add 30µL 1X lysis reagent to the assay wells and place the 96-well plate on an orbital shaker for one minute.
- 3. Remove plate from shaker and measure luminescence (as described in Section 13.4).

13.4 Measurement of Luminescence

- Luminescence is measured in the range of 300 to 650 nm, using an injecting luminometer and with
- software that controls the injection volume and measurement interval. Light emission from each well is
- expressed as relative light units (RLU) per well. The luminometer output is saved as raw data in an
- Excel® spread sheet. A hard copy of the luminometer raw data should be signed, dated and stored in the
- 953 study notebook.

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954 13.5 Data Analysis

- 955 BG1LUC4E2 ER TA uses an Excel® spreadsheet to collect and adjust the RLU values obtained from the
- 956 luminometer and a GraphPad Prism® template to analyze and graph data. Plate reduction is calculated
- 957 using unadjusted RLU values.
- The Excel® spreadsheet subtracts background luminescence (average DMSO solvent control RLU value)
- 959 from test substance, reference standard and control RLU values. Test substance, reference standard, and
- ontrol RLU values are then adjusted relative to the highest Ral/E2 reference standard RLU value, which
- is set to 10,000. After adjustment, values are transferred to GraphPad Prism® for data analysis and
- 962 graphing.

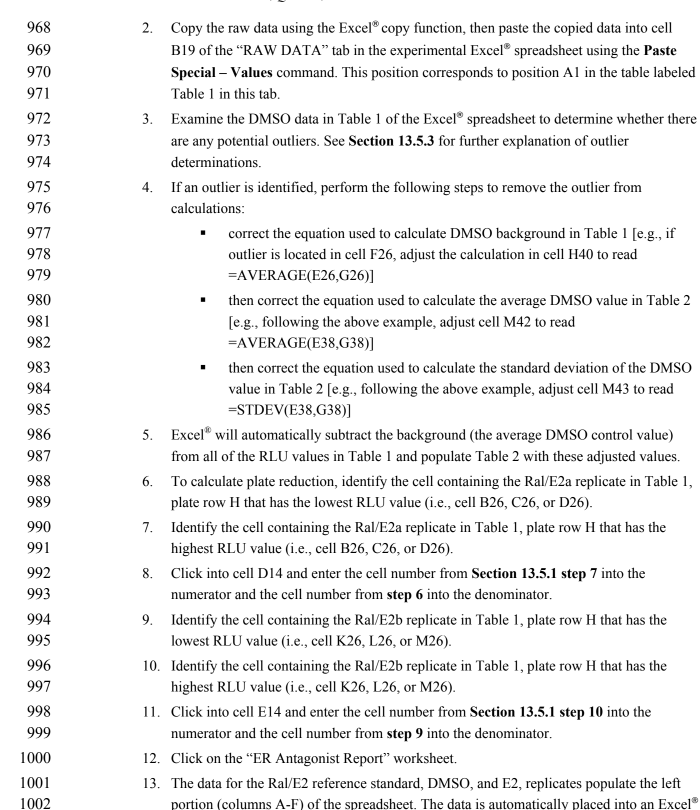
963 13.5.1 Collection and Adjustment of Luminometer Data for Range Finder Testing

- The following steps describe the procedures required to populate the Excel® spreadsheet that has been configured to collect and adjust the RLU values obtained from the luminometer.
- 966 1. Open the raw data file and the corresponding experimental Excel® spreadsheet from Section 13.1.1.

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graph.



14. To set the highest RLU value for the reference standard to 10,000 RLU, go to cell C2 of

"ER Antagonist Report" worksheet and check the formula contained within that cell. The

1006 1007		divisor should be the cell number of the cell containing the highest averaged Ral/E2 RLU value (column A).
1008 1009 1010	1	Open the "Visual Observation Scoring" worksheet. Enter the visual observation scores for each well on the 96-well plate. This data will be linked to the "ER Antagonist Report" worksheet.
1011 1012	1	6. After the testing results have been evaluated and reviewed for quality control, enter the following information into the Compound Tracking worksheet:
1013 1014		 Enter pass/fail results for plate reference standard and control parameters into the Plate Pass/Fail Table
1015 1016		 Enter information from the testing of coded substances into the Testing Results Table
1017 1018		■ Reviewer Name – Enter the name of the person who Reviewed\QC'ed the data into cell A34
1019		■ Date – Enter the date on which the data was reviewed into cell D34
1020	13.5.2 <u>C</u>	ollection and Adjustment of Luminometer Data for Comprehensive Testing
1021 1022 1023 1024		g steps describe the procedures required to populate the Excel [®] spreadsheet that has been collect and adjust the RLU values obtained from the luminometer. Open the raw data file and the corresponding experimental Excel [®] spreadsheet from Section 13.1.2.
1025 1026 1027 1028	2	Copy the raw data using the Excel® copy function, then paste the copied data into cell B14 of the "RAW DATA" tab in the experimental Excel® spreadsheet using the Paste Special – Values command. This position corresponds to position A1 in the table labeled Table 1 in this tab.
1029 1030 1031	3	Examine the DMSO data in Table 1 of the Excel® spreadsheet to determine whether there are any potential outliers. See Section 13.5.3 for further explanation of outlier determinations.
1032 1033	4	. If an outlier is identified, perform the following steps to remove the outlier from calculations:
1034 1035 1036		 correct the equation used to calculate DMSO background in Table 1[e.g., if outlier is located in cell M14, adjust the calculation in cell H40 to read =AVERAGE(M15:M17)]
1037 1038 1039		• then correct the equation used to calculate the average DMSO value in Table 2 [e.g., following the above example, adjust cell M35 to read =AVERAGE(M25:M27)]
1040 1041 1042		• then correct the equation used to calculate the standard deviation of the DMSO value in Table 2 [e.g., following the above example, adjust cell M36 to read =STDEV(M25:M27)]

1043 1044	5	Excel® will automatically subtract the background (the average DMSO control value) from all of the RLU values in Table 1 and populate Table 2 with these adjusted values.
1045 1046	6	To calculate plate reduction, identify the cell containing the Ral/E2 replicate in plate row G that has the lowest RLU value.
1047 1048	7	. Identify the cell containing the Ral/E2 replicate in plate row G that has the highest RLU value.
1049 1050	8	Click into cell D14 and enter the cell number from Section 13.5.2 step 7 into the numerator and the cell number from step 6 into the denominator.
1051 1052	9	. Identify the cell containing the Ral/E2 replicate in plate row H that has the lowest RLU value.
1053 1054	1	0. Identify the cell containing the Ral/E2 replicate in plate row H that has the highest RLU value.
1055 1056	1	1. Click into cell E14 and enter the cell number from Section 13.5.2 step 10 into the numerator and the cell number from step 9 into the denominator.
1057	1	2. Click on the "ER Antagonist Report" worksheet.
1058 1059 1060	1	3. The data for the Ral/E2 reference standard, DMSO, E2, and Flavone/E2 replicates populate the left portion (columns A-E) of the spreadsheet. The data is automatically placed into an Excel® graph.
1061 1062 1063 1064	1	4. To set the highest RLU value for the reference standard to 10,000 RLU, go to cell D2 of "ER Antagonist Report" worksheet and check the formula contained within that cell. The divisor should be the cell number of the cell containing the highest averaged Ral/E2 RLU value (column A).
1065 1066 1067	1	5. Open the "Visual Observation Scoring" worksheet. Enter the visual observation scores for each well on the 96-well plate. This data will be linked to the "ER Antagonist Report" worksheet.
1068 1069	1	6. Copy the data into GraphPad Prism® for the calculation of IC ₅₀ values and to graph experimental results as indicated in the NICEATM Prism® Users Guide.
1070 1071	1	7. After the testing results have been evaluated and reviewed for quality control, enter the following information into the Compound Tracking worksheet:
1072 1073		 Enter pass/fail results for plate reference standard and control parameters into the Plate Pass/Fail Table
1074 1075		 Enter information from the testing of coded substances into the Testing Results Table
1076 1077		 Reviewer Name – Enter the name of the person who Reviewed\QC'ed the data into cell A34
1078		 Date – Enter the date on which the data was reviewed into cell D32
1079	13.5.3 <u>I</u>	Determination of Outliers

The Study Director will use good statistical judgment for determining "unusable" wells that will be excluded from the data analysis and will provide an explanation in the study notebook for any excluded data. This judgment for data acceptance will include Q-test analysis.

1083 The formula for the Q test is:

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Outlier - Nearest Neighbor

Range (Highest – Lowest)

where the outlier is the value proposed for exclusion, the nearest neighbor is the value closest to the outlier, and the range is the range of the three values (Q values for samples sizes from 3 to 10 are provided in Table 13-2). For example, if the value of this ratio is greater than 0.94 (the Q value for the 90% confidence interval for a sample size of three) or 0.76 (the Q value for the 90% confidence interval for a sample size of four), the outlier may be excluded from data analysis.

Table 13-2 Q Test Values

Number Of Observations	Q Value
2	-
3	0.94
4	0.76
5	0.64
6	0.56
7	0.51
8	0.47
9	0.44
10	0.41

For E2 reference standard replicates (sample size of two), any adjusted RLU value for a replicate at a given concentration of E2 is considered and outlier if its value is more than 20% above or below the adjusted RLU value for that concentration in the historical database.

1094 13.5.4 <u>Acceptance Criteria</u>

1095 13.5.4.1 Range Finder Testing

Acceptance or rejection of a range finder test is based on reference standard and solvent control results from each experiment conducted on a 96-well plate.

- Reduction: Plate reduction, as measured by dividing the averaged highest Ral/E2
 reference standard RLU value by the averaged DMSO control RLU value, must be
 greater than three-fold.
- E2 control results: E2 control RLU values must be within 2.5 times the standard deviation of the historical E2 control mean RLU value (See Section 16.1).
- DMSO control results: DMSO control RLU values must be within 2.5 times the standard deviation of the historical solvent control mean RLU value (see **Section 16.2**).

An experiment that fails either acceptance criterion will be discarded and repeated.

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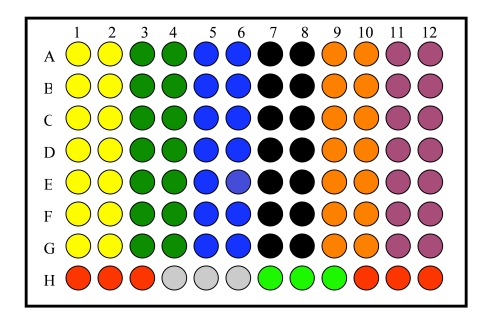
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1107	13.5.4.2 Comprehensive Testing
1108	Acceptance or rejection of a test is based on evaluation of reference standard and control results from
1109	each experiment conducted on a 96-well plate. Results are compared to quality controls (QC) for these
1110	parameters derived from the historical database (see Section 16.5), which are summarized below.
1111	 Reduction: Plate reduction, as measured by dividing the averaged highest Ral/E2
1112	reference standard RLU value by the averaged lowest Ral/E2 control RLU value, must be
1113	greater than three-fold.
1114	• DMSO control results: DMSO control RLU values must be within 2.5 times the standard
1115	deviation of the historical solvent control mean RLU value (see Section 16.5).
1116	• Reference standard results: The Ral\E2 reference standard concentration-response curve
1117	should be sigmoidal in shape and have at least three values within the linear portion of
1118	the concentration-response curve.
1119	• E2 control results: E2 control RLU values must be within 2.5 times the standard
1120	deviation of the historical E2 control mean RLU value.
1121	• Positive control results: Flavone/E2 control RLU values must be less than the E2 control
1122	mean minus three times the standard deviation from the E2 control mean.
1123	An experiment that fails any single acceptance criterion will be discarded and repeated.
1124	14.0 RANGE FINDER TESTING

RANGE FINDER TESTING

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1125 Antagonist range finding for coded substances consists of seven point, 1:10 serial dilutions tested in 1126 duplicate wells of the 96-well plate. Figure 14-1 contains a template for the plate layout used in antagonist range finder testing. 1127





- **DMSO** (Solvent Control)
- Range Finder for Sample #1
- Range Finder for Sample #2
- Range Finder for Sample #3
- Range Finder for Sample #4
- Range Finder for Sample #5
- Range Finder for Sample #6
- E2 Control

Figure 14-1 Antagonist Range Finder Plate Layout

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Evaluate whether range finder experiments have met acceptance criteria (see Section 13.6.3).

To determine starting concentrations for comprehensive testing use the following criteria:

• If results in the range finder test suggest that the test substance is negative for antagonist activity (i.e., if there are no points on the test substance concentration curve that are less than the mean minus three times the standard deviation of the E2 control, see **Figure 14-2**), comprehensive testing will be conducted using an 11-point 1:2 serial dilution using

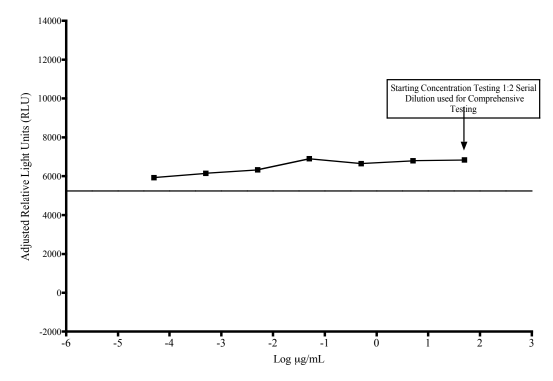
the maximum soluble concentration of test substance as the with the limit dose as the starting concentration.

- If results in the range finder test suggest that the test substance is negative for agonist activity (i.e., if there are no points on the test substance concentration curve that are greater than the mean plus three times the standard deviation of the DMSO control), and the higher concentrations in the range finder are cytotoxic, comprehensive testing will be conducted using an 11 point 1:2 serial dilution with the lowest cytotoxic concentration as the starting concentration (see **Figure 14-3**).
- If results in the range finder test suggest that the test substance is positive for antagonist activity (i.e., if there are points on the test substance concentration curve that are less than the mean minus three times the standard deviation of the E2 control), the top concentration to be used for the 11-point dilution scheme in comprehensive testing should be one of the following:
 - The concentration giving the lowest adjusted RLU value in the range finder
 - The maximum soluble concentration (See **Figure 14-2**)
 - The lowest cytotoxic concentration (See **Figure 14-3** for a related example).

The 11-point dilution scheme will be based on either a 1:2 or 1:5 serial or dilution according to the following criteria:

- An 11-point 1:2 serial dilution should be used if the resulting concentration range (note: an 11-point 1:2 serial dilution will cover a range of concentrations over approximately three orders of magnitude [three logs]) will encompass the full range of responses based on the concentration response curve generated in the range finder test (see Figure 14-4).
- If the concentration range that would be generated with the 1:2 serial dilution will not encompass the full range of responses based on the concentration response curve in the range finder test (see Figure 14-5), an 11-point 1:5 serial dilution should be used instead.
- If a substance exhibits a biphasic concentration response curve in the range finder test (see **Figure 14-6**), both phases should also be resolved in comprehensive testing. In this case, two peaks could potentially be used to identify the top concentration to be used for the 11-point dilution scheme in comprehensive testing. In order to resolve both curves, the top concentration should be based on the peak associated with the higher concentration and the top dose one log concentration higher than the concentration giving the lowest adjusted RLU value in the range finder. An 11-point 1:5 serial dilution should be used.

1171 Figure 14-2 Antagonist Range Finder (example 1)

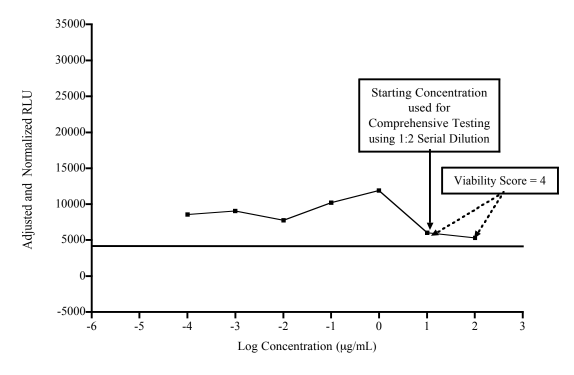


The solid horizontal line represents the mean minus three times the standard deviation of the E2 control.

1174 Figure 14-3 Antagonist Range Finder (example 2)

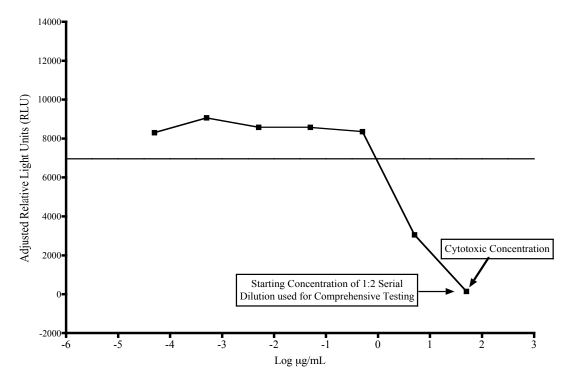
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1176 The solid horizontal line represents the mean minus three times the standard deviation of the E2 control.

1177 Figure 14-4 Antagonist Range Finder (example 3)

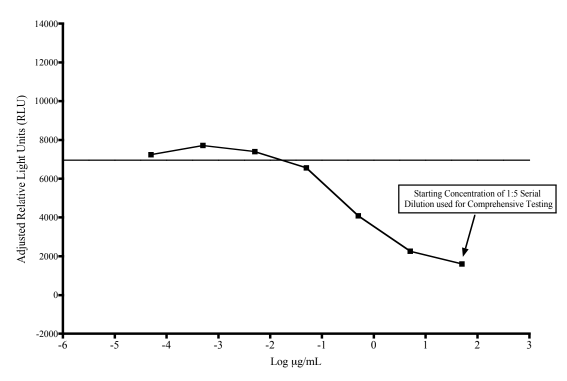


The solid horizontal line represents the mean minus three times the standard deviation of the E2 control.

1180 Figure 14-5 Antagonist Range Finder (example 4)

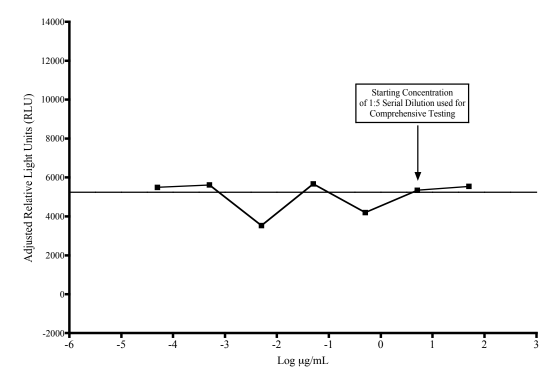
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The solid horizontal line represents the mean minus three times the standard deviation of the E2 control.

1183 Figure 14-6 Antagonist Range Finder (example 5)



The solid horizontal line represents the mean minus three times the standard deviation of the E2 control.

15.0 COMPREHENSIVE TESTING

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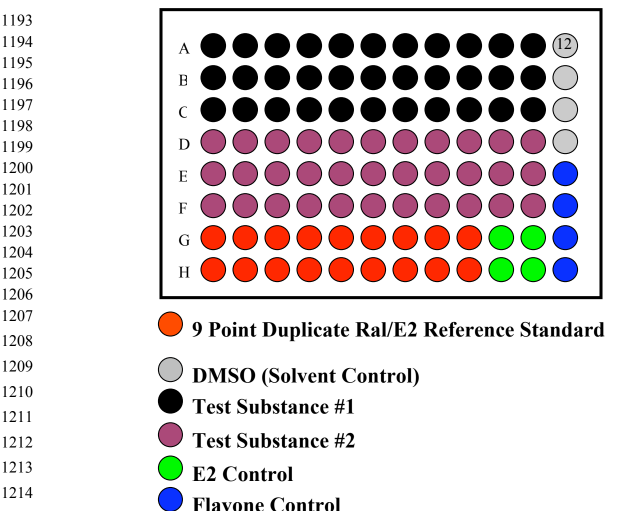
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1191 1192 Antagonist comprehensive testing for coded substances consists of 11 point, 1:2 serial dilutions, with each concentration tested in triplicate wells of the 96-well plate. **Figure 15-1** contains a template for the plate layout to be used in antagonist comprehensive testing.

Figure 15-1 Antagonist Comprehensive Test Plate Layout



Evaluate whether comprehensive experiments have met acceptance criteria (see **Section 13.6.3**) and graph the data as described in the NICEATM Prism® users guide.

- If the substance has been tested up to the limit dose or the maximum soluble dose without causing a significant decrease in cell viability, and there are no points on the concentration curve that are less than the mean minus three times the standard deviation of the E2 control, the substance is considered negative for antagonism.
- If the substance has been tested up to the limit dose and there are points on the concentration curve that are less than the mean minus three times the standard deviation of the E2 control, but cell viability has a visual inspection score of 2 or greater, at all points falling below the E2 line, the substance is considered negative for antagonism.
- If there are points on the test substance concentration curve that are less than the mean minus three times the standard deviation of the E2 control that do not cause a visual inspection score of 2 or greater, the substance is positive for antagonism.

1229 Points in the test substance concentration curve that cause a visual inspection 1230 score of 2 or greater, are not included in data analyses. 1231 COMPILATION OF THE HISTORICAL QUALITY CONTROL DATABASE 16.0 1232 Historical databases are maintained in order to ensure that the assay is functioning properly. Historical 1233 databases are compiled using Excel® spreadsheets and are separate from the spreadsheets used to collect 1234 the data for individual test plates. Reference standard and control data is used to develop and maintain the 1235 historical database and are used as quality controls to determine acceptance of individual test plates. 1236 The sources of data needed to compile the historical database for the E2 control and flavone/E2 control 1237 values are the experiment specific Excel® data collection and analysis spreadsheets (see Section 13.5.2) 1238 used for BG1LUC4E2 ER TA antagonist testing. The sources of the data needed to compile the historical 1239 database for the DMSO control are the experiment specific Excel® data collection and analysis 1240 spreadsheets used for BG1LUC4E2 ER TA antagonist and agonist testing (see Section 13.5.2 of the 1241 BG1LUC4E2 ER TA antagonist protocol and Section 11.5.2 in the BG1LUC4E2 ER TA agonist 1242 protocol). 1243 16.1 E2 Control 1244 Open the BG1LUC4E2 ER TA antagonist specific historical database Excel® spreadsheet 1245 (LUMI AgandAntQC.xls) and save under a new name using the Excel® "Save As" function, adding the 1246 laboratory designator to the file name (e.g., for Laboratory H, new name = HLUMI AgandAntQC.xls). 1247 Open the E2 Control worksheet and enter the date and experiment name into worksheet columns A and B 1248 respectively. Enter the experimental mean adjusted E2 control value (from cell D37 in the ER Antagonist 1249 Report worksheet of the Excel® data collection and analysis spreadsheet) into the Antagonist E2 control 1250 worksheet, column C. Acceptance or rejection of plate E2 control data for comprehensive testing is based 1251 on whether the mean plate E2 RLU value falls within 2.5 times the standard deviation of the E2 value in 1252 the historical database (columns G and H in the E2 Control worksheet). 1253 16.2 **DMSO** 1254 Open the combined agonist and antagonist BG1LUC4E2 ER TA historical database Excel® spreadsheet 1255 (LUMI AgandAntQC.xls) and save under a new name using the Excel® "Save As" function, adding the 1256 laboratory designator to the file name (e.g., for Laboratory H, new name = HLUMI AgandAntOC.xls). 1257 Enter the date and experiment name into worksheet columns A and B respectively. Enter the experimental 1258 mean DMSO control value (from cell H37 in the RAW DATA worksheet of the agonist and antagonist 1259 Excel® data collection and analysis spreadsheet) into worksheet column C. Acceptance or rejection of the 1260 plate DMSO control data for range finding and comprehensive testing is based on whether the mean plate 1261 DMSO RLU value falls within 2.5 times the standard deviation of the DMSO value in the historical 1262 database (columns G and H in the DMSO worksheet).

1263 17.0 QUALITY TESTING OF MATERIALS

- All information pertaining to the preparation and testing of media, media supplements, and other
- materials should be recorded in the Study Notebook.

1266 17.1 Tissue Culture Media

- Each lot of tissue culture medium must be tested in a single growth flask of cells before use in ongoing
- tissue culture or experimentation (**note:** each bottle within a given lot of Charcoal/Dextran treated FBS
- must be tested separately).

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- 1. Every new lot of media (RPMI and DMEM) and media components (FBS,
- 1271 Charcoal/Dextran treated FBS, and L-glutamine) must first be tested on the LUMI-
- 1272 CELL® ER assay prior to being used in any GLP acceptable assays.
- 1273 2. Add 4 μL of DMSO (previously tested) into four separate 13 mm tubes.
- 1274 3. Add 400 mL media (to be tested) to 13 mm tube.
 - 4. Dose an experimental plate as in **Section 12.0**, treating the media being tested as a test substance.
 - 5. Analyze 96-well plate as described in **Section 12.0**, comparing the data from the DMSO controls made using previously tested tissue culture media to the new media being tested.
 - 6. Use the agonist historical database to determine if the new media with DMSO lies within 2.5 standard deviations of the mean for the media. If the RLU values for the new media with DMSO lie within 2.5 standard deviations of the DMSO mean from the historical database, the new lot of media is acceptable. If the RLU values for the new media with DMSO do not lie within 2.5 standard deviations of the DMSO mean from the historical database, the new lot may not be used in the assay.
 - 7. Note date and lot number in study notebook.
 - 8. If the new bottle passes quality testing as described in **Section 15.1 step 6**, apply the media to a single flask cells and observe the cells growth and morphology over the following 2 to 3 days. If there is no change in growth or morphology, the new media is acceptable for use.

1290 **17.2 G418**

- 1. New lots of G418 must first be tested on the LUMI-CELL® ER assay prior to being used in any GLP acceptable assays.
 - 2. Add 220 μL of G418 (previously tested) to a single flask containing cells growing in RPMI.
 - 3. Add 220 µL of G418 (to be tested) to a different flask containing cells growing in RPMI.
- 1296 4. Observe cellular growth and morphology in both tissue culture flasks over a 48 to 72
 1297 hour period. If there are no differences in observed growth rate and morphology between
 1298 the two flasks, the new G418 lot is acceptable.

1300 G418 is not acceptable. 1301 Note date and lot number in study book. 1302 **DMSO** 17.3 1303 Every new bottle of DMSO must be tested on the LUMI-CELL® ER assay prior to use in 1304 any GLP acceptable assays. 1305 2. Add 4 µL of DMSO (to be tested) into four separate 13 mm tubes. 1306 3. Add 400 mL media (previously tested) the same tubes. 1307 4. Dose an experimental plate as in Section 15.0, treating the media being tested as a test 1308 substance. 1309 Analyze 96-well plate as described in **Section 15.0**, comparing the data from the DMSO 1310 controls made using previously tested tissue culture media to the new media being tested. 1311 Use the agonist historical database to determine if media with new DMSO lies within 2.5 1312 standard deviations of the DMSO mean from historical database. If the RLU values for 1313 the media with new DMSO lie within 2.5 standard deviations of the DMSO mean from 1314 the historical database, the new lot of DMSO is acceptable. If the RLU values for media 1315 with new DMSO do not lie within 2.5 standard deviations of the DMSO mean from 1316 historical database, the new lot may not be used in the assay. 1317 Note the date, lot number, and bottle number in study book. 1318 If no DMSO has been previously tested, test several bottles as described in Section 15.3, 1319 and determine whether any of the bottles of DMSO have a higher average RLU than the 1320 other bottle(s) tested. Use the DMSO with the lowest average RLU for official 1321 experiments. 1322 17.4 **Plastic Tissue Culture Materials** 1323 Grow one set of cells, plate them for experiments on plastic ware from the new lot and 1324 one set of cells in the plastic ware from a previous lot, and dose them with E2 reference 1325 standard and controls. 1326 Perform the BG1LUC4E2 ER TA experiment with both sets of cells. 1327 If all of the analysis falls within acceptable QC criteria, then the new manufacturer's 1328 products may be used. 1329 Eli Lilly and Company and National Institutes of Health Chemical Genomics Center. 2005. Assay 1330 Guidance Manual Version 4.1. Bethesda, MD: National Institutes of Health. Available: 1331 http://www.ncgc.nih.gov/guidance/manual_toc.html [accessed 05 September 2006] 1332 ICCVAM. 2001. Guidance Document on Using In Vitro Data to Estimate In Vivo Starting Doses for 1333 Acute Toxicity. NIH Pub. No. 01-4500. Research Triangle Park, NC: National Institute of Environmental

If cellular growth is decreased, or the cells exhibit abnormal morphology, the new lot of

- Health Sciences. Available: http://iccvam.niehs.nih.gov/methods/invidocs/
- guidance/iv_guide.pdf [accessed 31 August 2006]